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The frequency and clinical presentation of Zika virus coinfections: a systematic review

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ABSTRACT

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Correspondence to Dr Elizabeth B Brickley; elizabeth.brickley@lshtm.ac.uk **Background** There is limited knowledge on the influence of concurrent coinfections on the clinical presentation of Zika virus (ZIKV) disease.

Methods To better understand the types, frequencies and clinical manifestations of ZIKV coinfections, we did a systematic review of four databases (PubMed, Embase, Web of Science, LILACS) without restrictions for studies on ZIKV coinfections confirmed by nucleic acid (quantitative real-time-PCR) testing of ZIKV and coinfecting pathogens. The review aimed to identify cohort, cross-sectional, case series and case report studies that described frequencies and/or clinical signs and symptoms of ZIKV coinfections. Conference abstracts, reviews, commentaries and studies with imprecise pathogen diagnoses and/or no clinical evaluations were excluded.

Results The search identified 34 articles from 10 countries, comprising 2 cohort, 10 cross-sectional, 8 case series and 14 case report studies. Coinfections were most frequently reported to have occurred with other arthropod-borne viruses (arboviruses); out of the 213 coinfections described, ZIKV infections co-occurred with chikungunya in 115 cases, with dengue in 68 cases and with both viruses in 19 cases. Other coinfecting agents included human immunodeficiency, Epstein-Barr, human herpes and Mayaro viruses, *Leptospira* spp, *Toxoplasma gondii* and *Schistosoma mansoni*. ZIKV-coinfected cases primarily presented with mild clinical features, typical of ZIKV monoinfection; however, 9% of cases in cohort and cross-sectional studies were reported to experience complications.

Conclusion Based on the evidence collated in this review, coinfections do not appear to strongly influence the clinical manifestations of uncomplicated ZIKV infections. Further research is needed to confirm whether risk of severe complications is altered when ZIKV infection co-occurs with other infections.

PROSPERO registration number CRD42018111023.

INTRODUCTION

Zika virus (ZIKV) is an *Aedes* mosquito-borne flavivirus that recently emerged in the Americas.¹ First recognised in Brazil in early 2015,

Key questions

What is already known?

As Zika virus (ZIKV) has been most prevalent in subtropical and tropical regions with high burdens of cocirculating infectious agents, a proportion of ZIKV infections occur simultaneously with infections by one or multiple other pathogens; however, it is uncertain whether coinfections may influence ZIKVrelated pathology.

What are the new findings?

- This systematic review collated the evidence on ZIKV coinfections as published in 34 studies in 10 countries. ZIKV coinfections were most frequently reported in the context of the arthropod-borne viruses, dengue and chikungunya, but were also described in relation to eight other pathogens.
- While the findings of this review suggest that coinfections do not appear to strongly influence the clinical manifestations of uncomplicated ZIKV infections, this review did identify reports of neurological complications in the context of coinfection.

What do the new findings imply?

The findings of this review highlight a need for coordinated and rapid research efforts during future outbreaks to optimise diagnostic testing strategies for detecting coinfections and determining whether they may exacerbate the risk of severe ZIKV complications, such as Guillain-Barré syndrome and congenital Zika syndrome.

the ZIKV epidemic spread explosively, with autochthonous transmission reported in more than 86 countries and territories by 2018.¹ Given the widespread circulation of this emerging infection of public health concern, it is critical that healthcare practitioners can readily recognise ZIKV disease across the full range of its clinical presentations.

Current evidence indicates that ZIKV infections typically present with no or mild clinical features.¹ A 2018 meta-analysis of 23 studies

by Haby and colleagues estimated a prevalence of asymptomatic ZIKV infections of 62% (95% CI 33% to 87%).² For symptomatic ZIKV disease, the WHO describes a mild clinical presentation marked by fever, rash, conjunctivitis, myalgia, arthralgia, malaise and headache.¹ Nevertheless, ZIKV is neurotropic and, in a subset of cases, infections have been associated with severe neurological complications, including the polyneuropathy Guillain-Barré syndrome (GBS) and congenital Zika syndrome (CZS), a constellation of congenital central nervous system malformations resulting from the vertical transmission of ZIKV during pregnancy.³ It has been estimated that GBS arises in approximately 2 per 10 000 ZIKV infections,¹⁴ and the absolute risk of adverse birth outcomes (ie, miscarriage, stillbirth, premature birth and CZS) has been reported to range between 7% and 46% in pregnancies with quantitative real-time PCR (qRT-PCR)-confirmed ZIKV infection.5-8

Although the clinical presentation of ZIKV monoinfections has been well characterised, one factor that may influence the clinical spectrum of ZIKV disease is coinfection. Given the high incidence of infectious diseases in the subtropical and tropical areas where ZIKV is prevalent, a proportion of all ZIKV infections occur concurrently with infections by one or multiple pathogens.⁹ ZIKV disease in the context of coinfection remains inadequately investigated, and it is uncertain whether specific coinfections may influence the presentation and severity of ZIKV-related signs and symptoms. A 2019 literature review by Vogels and colleagues hypothesised that coinfecting agents have the potential to enhance, inhibit, compete with or have no effect on ZIKV replication and the resulting clinical disease.¹⁰ To advance understanding on this topic, this systematic review aims to quantify how frequently ZIKV coinfections occur among ZIKV-infected populations and to investigate whether the clinical course of ZIKV disease in humans is altered in the context of coinfection.

METHODS

Search

Four databases (PubMed, Web of Science, LILACs and EMBASE) were searched for publications up to 19 October 2019 using a comprehensive search strategy (online supplementary appendix 1). Keywords and Medical Subject Headings linked to ZIKV, bacterial, parasitic and other viral infectious diseases were used. The search included English, French, Spanish and Portuguese terms. No date or language restrictions were applied. The systematic review was registered in PROS-PERO. All study titles and abstracts were screened based on eligibility criteria, and references of included studies were also screened to identify additional eligible articles.

Study selection and data extraction

Cohort studies, cross-sectional studies, case series and case reports describing coinfections of ZIKV with one

or multiple other pathogens, confirmed by nucleic acid testing (eg, qRT-PCR) for ZIKV, and all coinfecting pathogens were eligible for inclusion in the review. Recovery of live pathogens was also considered to be indicative of acute coinfection. Of note, HIV-positive ZIKV cases with HIV suppression were not included in this review. Two reviewers (AR and LL) simultaneously screened studies for eligibility, and any discrepancies were resolved by a third reviewer (EBB). Conference abstracts, reviews, commentaries and studies without nucleic acid confirmation were excluded. Whereas cohort, cross-sectional and case series studies reporting on numbers of ZIKV coinfections without description of signs and symptoms were included to describe the frequency of ZIKV coinfections, studies with no reporting of signs and symptoms of ZIKV coinfections were otherwise excluded from the review. Data extraction was independently performed by two reviewers (AR, LL). From the full-text articles, information on study author, location, year, data source, age and sex of identified cases was extracted. Additional extracted information included frequencies of ZIKV cases with coinfection, types of coinfection, types of diagnostic testing, reported signs and symptoms, noninfectious comorbidities, and types and frequencies of complications. To investigate the frequency of ZIKV coinfections in cohort, cross-sectional and case series studies, the numbers of coinfections out of the total number of qRT-PCR-confirmed ZIKV cases were calculated for the eligible studies. The study quality assessment was conducted using the Oxford Centre for Evidence-based Medicine (OCEBM) Levels of Evidence, March 2009¹¹; see online supplementary appendix 2 for details.

Patient and public involvement

This research was done without patient or public involvement.

RESULTS

Study selection

The search initially identified 12 253 titles, of which 12 050 titles were excluded after screening titles and abstracts and removing duplicates (figure 1). Full-text screening was completed for 203 publications, and, ultimately, 34 articles representing coinfections in 10 countries were included (tables 1–4 and figure 2).

ZIKV coinfection types

ZIKV infections were most frequently reported to occur concurrently with other arthropod-borne viruses (arboviruses). Out of the 213 coinfections examined, there were 115 ZIKV/chikungunya virus (CHIKV) coinfection cases, 68 ZIKV/dengue virus (DENV) coinfection cases and 19 cases coinfected with all three viruses. Other reported ZIKV coinfections included ZIKV/HIV (n=3), ZIKV/Leptospira spp (n=2), ZIKV/CHIKV/HIV/Toxoplasma gondii (n=1), ZIKV/CHIKV/Toxoplasma gondii (n=1), ZIKV/Epstein-Barr virus (EBV)/human herpes viruses-6 (HHV-6) (n=1), ZIKV/herpes simplex virus-1

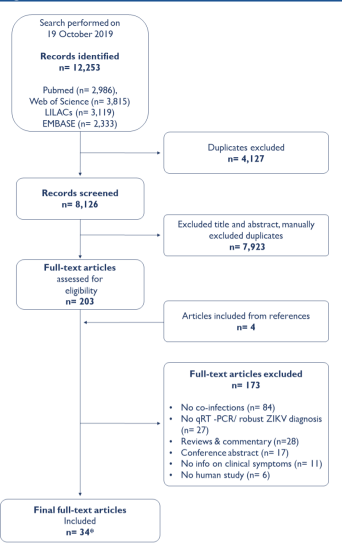


Figure 1 Study selection.

*Studies reporting on both clinical signs and symptoms and frequency of Zika virus coinfections (n=27); studies reporting only on Zika virus coinfection frequencies (n=7). qRT-PCR, quantitative real-time PCR; ZIKV, Zika virus.

(HSV-1) (n=1), ZIKV/Mayaro virus (MAYV) (n=1) and ZIKV/*Schistosoma mansoni* (n=1) (figure 3).

Frequencies of ZIKV coinfections

The frequencies of coinfections among ZIKV-infected populations were reported in 11 studies, including 1 cohort study, 7 cross-sectional studies and 3 case series (table 1, online supplementary table 2). Frequency estimates were reported only for coinfections with CHIKV and DENV and varied geographically and across study populations at risk. Among patients presenting with arbovirus-like symptoms, ZIKV/CHIKV coinfection frequencies were reported to range from 0.3% in a study in Colombia to 54% in a study in Brazil.^{5 9 12-14} Similarly, ZIKV/DENV coinfection frequencies in patients with arbovirus-like symptoms were reported to range from 0.03% in a study in Colombia to 47.4% in a study in Brazil.^{9 13 15-19} ZIKV/CHIKV/DENV coinfection frequencies ranged from 8% in a study in Nicaragua to 27.6% in a study in Colombia.^{9 13}

Signs and symptoms of coinfections

In total, 27 studies, including 1 cohort study, 5 crosssectional studies, 7 case series and 14 case report studies, reported the signs and symptoms of ZIKV coinfection across a total of 106 ZIKV-coinfected cases.

ZIKV/CHIKV coinfections

The clinical presentations of 48 cases with ZIKV/CHIKV coinfection were reported in 1 cohort study, 1 cross-sectional study, 4 case series and 6 case reports (tables 2–4, online supplementary tables 1 and 2).^{20–28} Within the cohort, cross-sectional and case series studies, cases were reported to present with the following signs and symptoms consistent with the WHO ZIKV clinical case definition¹: fever (33%–100%), rash (0%–100%), conjunctivitis (0%–50%), myalgia (67%–100%), arthralgia (0%–67%) and headache (17%–50%) (tables 2 and 3). In addition, gastrointestinal (GI) symptoms were reported in 17% to 100% of cases in three studies (tables 2 and 3).^{20 21 23}

Complications were reported among 14.7% (5 cases) of ZIKV/CHIKV-coinfected cases in cohort and crosssectional studies,^{22–27} of whom two adult cases presented with unspecified neurological complications that resulted in death (figure 4).²⁸ Additionally, two coinfections in pregnancy were associated respectively with an encephaly and an absence of a heartbeat.²⁸ A non-neurological complication reported was a case that died from multiorgan failure following haemorrhagic manifestations.^{23 28} The case series studies described that six out of eight ZIKV/CHIKV-coinfected cases developed complications, which included neurological manifestations, such as GBS in two cases,^{22 29} encephalitis in one case,²² myeloradiculitis in one case,²⁹ as well as non-neurological complications, such as persistent severe arthralgia in one case.²³ Additionally, four case reports described ZIKV/CHIKV coinfection-associated complications, including GBS in two cases,^{24 27} persistent severe arthralgia in one case²⁶ and sepsis resulting in death in one case.²

ZIKV/DENV coinfections

The clinical features of 42 cases with ZIKV/DENV coinfection were described across four cross-sectional studies, three case series and five case reports (tables 2–4, online supplementary tables 1 and 2).^{15 18 19 30–33} Cases with ZIKV/DENV coinfection within the cross-sectional and case series studies were reported to present with the following signs and symptoms consistent with the WHO ZIKV clinical case definition¹: fever (58%–100%), rash (53%–100%), conjunctivitis (25%–100%), myalgia (75%–100%), arthralgia (50%–100%) and headache (50%–100%) (tables 2 and 3). Other reported clinical features included GI symptoms in 17%–75% of cases and upper respiratory tract (URT) symptoms in 13%–25% of cases (tables 2 and 3).

Complications were reported among none of the ZIKV/DENV-coinfected individuals in cohort and cross-sectional studies (figure 4). However, seven cases with complications were reported in case series, which

ZIKV/CHIKV, ZIKV/DENV and ZIKV/CHIKV/DENV coinfection frequencies among qRT-PCR-confirmed ZIKV infected study population (n=11 studies) qRT-PCR-	IKV/CHIKV/DENV coinfection frequencie	infection frequencie	cie	s among qRT-PCR-confirm	ed ZIKV infe	cted study p	opulation (n=1 qRT-PCR-	1 studies)	
Country/ Territory Region Study year Study design Stu	Study year Study design	design	Stu	Study population*	Coinfecting agent(s)	Coinfection cases (n)	confirmed ZIKV infected study population (n)	Frequency (%)	Level of evidence†
Colombia N/A‡ Oct 2015- Cross sectional Susp Dec 2016	Oct 2015- Cross sectional Dec 2016	sectional	Susp	Suspected arbovirus infections	CHIKV	28	10118	0.3%	2c
Brazil South-East Sept 2015- Cohort Pregn May 2016	Sept 2015- Cohort May 2016		Pregn	Pregnant women with rash	CHIKV	с	182	1.7%	2b
Brazil North-East May 2015- Cross sectional Suspe May 2016	May 2015- Cross sectional May 2016	sectional	Suspe	Suspected arbovirus infections	CHIKV	٥	26	7.7%	2c
Nicaragua N/A Sept 2015- Cross sectional Suspec Apr 2016	Cross sectional	sectional	Suspec	Suspected arbovirus infections	CHIKV	16	75	21.3%	2c
Colombia East Aug 2015- Cross sectional Suspec Apr 2016	Cross sectional	sectional	Suspec	Suspected arbovirus infections	CHIKV	10	29	27.6%	2c
Brazil North-East Mar 2016– Cross sectional Suspecte May 2016 with rash	Mar 2016- Cross sectional May 2016	sectional	Suspec with ra	Suspected arbovirus infections with rash	CHIKV	36	66	54.0%	2c
Colombia N/A‡ Oct 2015- Cross sectional Suspec Dec 2016	Oct 2015- Cross sectional Dec 2016	sectional	Suspec	Suspected arbovirus infections	DENV	က	10118	0.03%	2c
Singapore Singapore Aug 2016– Case series Suspec Sept 2016	Aug 2016- Case series Sept 2016	series	Suspec	Suspected ZIKV infections	DENV	4	163	2.4%	4
Brazil North-East May 2015 Case series Suspec	May 2015 Case series	series	Suspec	Suspected arbovirus infections	DENV	-	31	3.2%	4
Brazil South-East Jan 2016- Cross sectional Susper Nov 2016	Jan 2016- Cross sectional Nov 2016	sectional	Suspec	Suspected ZIKV infections	DENV	4	100	4.0%	2c
Brazil South-East Jan 2016- Case series Susper Nov 2016	Jan 2016- Case series Nov 2016	series	Suspe	Suspected arbovirus infections	DENV	12	151	7.9%	4
5 - Cross sectional	Cross sectional	sectional	Suspec	Suspected arbovirus infections	DENV	9	75	8.0%	2c

Magalhaes *et al*'^{iz}

(2017)

Mercado-Reyes et al²⁸

Author (year)

Table 1

Brasil *et al*⁵

(2018)

(2016)

Waggoner et al¹³ (2016)

Carrillo- Hernández *et al*⁹

(2018)

Charlys da Costa *et al*¹⁴

(2017)

Mercado-Reyes *et al²⁸*

Case series	Suspected ZIKV infections	DENV	4	163	2.4%
Case series	Suspected arbovirus infections	DENV	-	31	3.2%
Cross sectional	Suspected ZIKV infections	DENV	4	100	4.0%
Case series	Suspected arbovirus infections	DENV	12	151	7.9%
Cross sectional	Suspected arbovirus infections	DENV	9	75	8.0%

2c

34.4%

29

42

DENV

Suspected arbovirus infections

Cross sectional

Aug 2015– Apr 2016

East

Colombia

Carrillo- Hernández *et al⁹*

(2018)

Estofolete *et al*¹⁸

(2018)

Colombo et al¹⁷

(2017)

Pessôa *et al*¹⁶

(2016)

Chia et al¹⁵

(2017)

(2018)

Waggoner *et al* (2016)¹³

Apr 2016

20

47.7%

38

8

DENV

Suspected ZIKV infections

Cross sectional

Feb 2016-

Central-West

Brazil

Mar 2016

2c

8.0%

75

20

27.6%

29

9	ω
CHIKV/ DENV	CHIKV/ DENV
Suspected arbovirus infections	Suspected arbovirus infections
Cross sectional	Cross sectional
Sept 2015– Apr 2016	Aug 2015– Apr 2016
N/A	East
Nicaragua	Colombia
Waggoner et al ¹³ (2016)	Carrillo-Hernández et a/ ⁹ (2018)

*Online supplementary table 2 gives details of all study populations in cohort studies and case reports.

TAll articles were rated according to level of evidence using the Oxford Centre for Evidence-based Medicine's Levels of Evidence, March 2009.¹¹ #Cases originated from the National Surveillance System in Public Health from Colombia. Therefore, cases come from all over Colombia, with the condition of living in a place 2200 m above sea level. CHIKV, chikungunya virus; DENV, dengue virus; N/A, not available; ZIKV, Zika virus.

Azeredo *et al¹⁹*

(2018)

		Level of evidence†								VEEV,
			2C	2b	2c	2c	2c	2c	20	tory tract;
	r of other igns and t (%)	GI symptoms	RN	17%	40%¶	RN	100%	RN	* HN	upper respirat
lies)	Frequency of other reported signs and symptoms (%)	URT symptoms	RN	RN	13%	RN	R	RN	**	tis virus; URT,
n=6 stuc		Headache	Щ	17%	67%	RN	100%	RN	*# 2	ouis encephali
ections (toms	Myalgia	RN	67%	93%	R	R	R	* 2	SLEV, Saint Lo
porting on signs and symptoms of qRT-PCR-confirmed ZIKV coinfections (n=6 studies)	Frequency of WHO ZIKV signs and symptoms $(\%)$	Conjunctivitis	RN	NR	60%	NR	RN	NR	*****	URT symptoms: pharyngits, sore throat, cough, pharyngeal congestion, adenopathy. GI symptoms: nausea, diarnhoea, vomiting, constipation, stomach ache. Online supplementary table 2 gives details of all study populations. Online supplementary table 2 gives details of all study populations. Online supplementary table 2 gives details of all study populations. Selevanteen asses of DENV-1 and one cases of DENV-1 and one cases with vomiting. Segens and symptoms were only reported for 15 patients. Teaves was accorded to r15 patients. Teaves was eached to no present with typical ZIKV signs but specific details were not reported. AFL acute februie liness: CHIKV chikungmya virus; DENV, dengue virus; EEV, east equine ercephalits virus; LHV, lineus virus; NAV, Mayaro virus; NR, not reported; ROCV, Rocio virus; SLEV, Saint Louis encephalits virus; URT, upper respiratory tract; VEEV,
onfirmed	VHO ZIKV s	Arthralgia	Ч	67%	87%	RN	100%	R	*HNN	ot reported; R(
CR-c	ncy of V	Fever	ЯN	33%	73%	ЯN		100%	NR**	us; NR, n
qRT-F	Freque (%)	Rash	НN	0	53%	R	100% NR	R	** NB	he. Mayaro vit
ptoms of	Other pathogens.		DENV	DENV2/ MAYV	CHIKV	CHIKV	CHIKV	RN	DENV, CHIKV, YFV, SLEV, ILHV, ROCV, WNV, WEEV, WEEV, VEEV	on, stomach ac 009. ¹¹ is virus; MAYV,
is and sym	z		28	9	18§	m		-	-	URT symptoms: pharynglits, sore throat, cough, pharyngeal congestion, adenopathy. GI symptoms: nausea, diarrhoea, vomiting, constipation, stomach ache. Colline supplementary table 2 gives details of all study populations. T-MI and each each of the lowest of the ordence using the Oxford Centre for Evidence-based Medicine's Levels of Evidence, March 2009. ¹¹ Esteventeen cases of DENV-1 and one case of DENV.4. Signs and symptoms were only reported for 15 patients. "For the cent of cases present of the apple and and cases with vomiting. "Case was described to present with typical ZIIV signs and 13% of cases with vomiting. ARI, and the cent of cases present with subcal 20% of cases with vomiting. ARI, and the cent of cases present with subcal ZIIV signs, and 13% of cases with vomiting. ARI, and the cent of cases present with subcal ZIIV signs. Devidence, the store the control for the cent of cases control and the case of cases with vomiting.
g on sigr		Coinfecting agent(s)	CHIKV	CHIKV	DENV‡	DENV	DENV	DENV	MAYV	i, diarrhoea, vo s's Levels of Ev s, GI, gastrointe
s reportin	N. total	study Coinfect population agent(s)	23 871	252	134	23 871	თ	139	453	otoms: nausea ased Medicine cot reported. contaitis virus
studies	Mean		58	7.5	34	28	Щ	RN	원	ny. Gl symr :vidence-b ails were r equine end
ional s		% female	74	48	ЯN	74	ЧN	63	20	adenopath entre for E vomiting. pecific dei EEV, east
Summary of cohort and cross-sectional studies rel		Study population*	Suspected arbovirus infections	AFI cases	Suspected arbovirus infections	Suspected arbovirus infections	AFI cases	AFI cases	AFI cases	URT symptoms: pharyngits, sore throat, cough, pharyngaal congestion, adenopathy. Gl symptoms: naus Colline supplementary table 2 gives details of all study populations. T-M and the supplementary table 2 gives details of all study populations. Edeventeen cases of DBN-1 and one case of DENV-4. SSigns and symptoms were only reported for 15 patients. "Case was described to presented with nauses. IDENV digoup virus; EFOX dependencies details were not reported. "Case was described to present with typical ZIAN signs for 3 days, but specific details were ond reported. ARI, and concerning to an onconvention of the table and onconcerning the concerning of the case of the typical ZIAN signs to a concerning the concerning the reported.
ort and c		Study Study Study	Cross- sectional study	Cohort study	Cross- sectional a study ii	Cross- sectional study	Cross- sectional study	Cross- / sectional study	Cross- sectional study	ugh, pharyng, of all study pr f evidence usil o of DENV-4. or 15 patients. usea, and 139 usea, and 137 ya virus; DENV
ry of coh		Study year	Colombia 2015/2016	2014/2015	2016	Colombia 2015/2016	2014/2016	2016	2015/2016	URT symptoms: pharyngtis, sore throat, cough, pharyngeal congest "Online supplementary table 2 gives details of all study populations. "Thal anciens were rated according to level of whoteno using the Oxt 456 worteen cases of DENV-1 and one case of DENV-4. "Signs and symptoms were only reported for 15 patients. "Case was described to present with typical ZINK signs for 3 days," "Case was described to present with typical ZINK signs for 3 days."
Summa		Stud Location year	Colombia	Haiti	Brazil	Colombia	Brazil	Реп	Brazil	s: pharyngitis, imentary table are rated acco. ises of DENV- mptoms were t of cases pre: is cribed to pre is cribed to pre is cribed to pre- is cribed to pre-
Table 2		Author (year)	Mercado- Reyes <i>et</i> al ²⁸ (2018)	Ball <i>et al²⁰</i> (2018)	Azeredo <i>et</i> al ¹⁹ (2018)	Mercado- Reyes <i>et</i> a/ ²⁸ (2018)	Araújo <i>et</i> a/ ³⁸ (2019)	Alva-Urcia et al ³⁰ (2016)	de Souza Costa ef al ⁴² (2019)	URT symptom *Online supple †All atricles we ‡Seventeen ca \$Signs and sy ¶Forty per can *Case was de AFI, acute febr Venezitalen ent

Table 3	Summe	ary of case	series	Summary of case series studies reporting on signs	orting	on sign	is and syr	nptoms c	of differen	t qRT-PCF	R-confi	rmed Z	ZIKV co	and symptoms of different qRT-PCR-confirmed ZIKV coinfections (n=7 studies)	n=7 stu	dies)			
						Mean	N, total		z	Other	Frequer (%)	icy of WH	HO ZIKV s	Frequency of WHO ZIKV signs and symptoms $\left(^{\phi _{0}} ight)$	toms		Frequency of other reported signs and symptoms (%)	of other gns and (%)	
Author (year)	Location	Study Study year design	Study design	Study population*	% female	age (years)	idy pulation	Coinfecting agent(s)			Rash F	Fever A	thralgia	Arthralgia Conjunctivitis	Myalgia	Headache	URT symptoms	GI symptoms	Level of evidence†
Acevedo <i>et</i> al (2017) ²²	Ecuador	2016	Case series	Cases with neurological symptoms‡	38	42	16	CHIKV	ო	Ś	E E E	100% 33	33%	R	RN	33%	R	RN	4
Metha <i>et al</i> (2018) ²⁹	Brazil	2015/2016	Case series	Cases with neurological symptoms§	50	52	22	CHIKV	5	DENV	100%	50% 50	50%	RN	R	NR	NR	NR	4
Sardi <i>et al</i> (2016) ²³	Brazil	2015	Case series	AVI and of qRT- NR PCR ZIKV+ infections	RN	RN	15	CHIKV	5	DENV	50% 1	100% 50	50%	50%	100%	50%	RN	50%	4
Cabral- Castro <i>et al</i> (2016) ²¹	Brazil	2015/2016	Case series	Suspected DENV infections	R	NR	30	CHIKV		DENV	100% 100%	0 %00		0	RN	NR	RN	100%	4
Estofolete <i>et</i> a/ (2018) ¹⁸	et Brazil	2016	Case series	Suspected arbovirus infections	42	46	1254	DENV	12	CHIKV	58%	58% 50	50%	25%	83%	75%	RN	17%	4
Chia <i>et al</i> (201 <i>T</i>) ¹⁵	Singapore	e 2016	Case series	Suspected ZIKV infections	RN	NR	163	DENV	4	NR	100%	100% 50	50%	50%	75%	50%	25%	75%	4
Li <i>et al</i> (2017) ³¹	Singapore	e 2016	Case series	Suspected ZIKV infections	50	11	14	DENV		CHIKV	100%	100% 10	100%	100%	100%	100%	0	RN	4
Acevedo <i>et</i> al (2017) ²²	Ecuador	2015/2016	Case series	Cases with neurological symptoms‡	38	42	9	CHIKV, DENV	4	Ś	RN	50% NR		ЧZ	RN	25%	25%	25%	4
Acevedo et al (2017) ²²	Ecuador	2016	Case series	Cases with neurological symptoms‡	33	42	9	CHIKV, HIV, Toxo		Ś	R	100% NR		R	R	100%	R	RN	4
URT sympton *Online supple +All articles w	ns: pharyngitis ementary table ere rated acco	URT symptoms: pharyngitis, sore throat, cough, pharyngeal conges of Online supplementary table 2 gives details of all study populations. t-MI articles wave rated according th level of evidence using the DVr	ugh, pharyi of all study	URT symptoms: pharyngitis, sore throat, cough, pharyngeal congestion, adenopathy. Gl symptoms: Online supplementary table 2 gives details of all study populations. #All articles ware rated according to level of exidence using the Oxford Centre for Evidence-based I	denopathy. ntre for Fvic	. Gl symptor		: nausea, diarrhoea, vomiting, constipation, s Medicine's Levels of Evidence March 2009 ¹¹	g, constipation,	nausea, diarrhoea, vomiting, constipation, stomach ache. Aedicine's I evals of Evidence March 2009 ¹¹									

1 at laticles were rated according to level of evidence using the Oxford Centre for Evidence-based Medicine's Levels of Evidence, March 2009¹¹ Edesociated with susperdent arbohrust infection. Stested for DENN; anstant, HSV/IXE, GMK, GMK, ERV, VZ, Toxo, MTB, enterovirus. AVI, acute viral illness: CHIKV, chikungunya virus; CMV, cytomegalovirus; DENV, dengue virus; GH, gastrointestinat, HSV, hence simplex virus; MT, Mycobacterium tuberculosis ; NR, not reported; Toxo, Toxoplasma gondii, URT, upper respiratory tract; VZ, varicella zoster.

Lobkowicz L, et al. BMJ Global Health 2020;5:e002350. doi:10.1136/bmjgh-2020-002350

Summary of case reports reporting on signs and symptoms of qRT-PCR-confirmed ZIKV coinfections (n=21 reports)

Table 4

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						Other	Z OHW	IKV sign	WHO ZIKV signs or symptoms	SU			Other reported signs or symptoms	ted signs or		
Author (year)	Location	Study year	A Sex ()	Age ((years) a	Age Coinfecting Sex (years) agent(s)	tested (negative)	Rash	Fever	Arthralgia (Conjunctivitis	Myalgia	Myalgia Headache	URT symptoms	GI symptoms	Additional information	Level of evidence*
Brito <i>et al²⁴</i> (2017)	Brazil	2016	× W	74 0	CHIKV	DENV, HIV, CMV, HTLV, Schisto, HSVI/2 cystcercosis	RN	+	+	E N	Ч	К	R	÷	Complications: Meningoencephalitis associated with Guillain- Barré syndrome (EMG confirmed)	υ
Silva et al (2018)	Brazil	2016	ε Σ	30	CHIKA	DENV, bacterial/fungal infections	۳	+	+	ц.	+	щ	щ	٣	Comorbidities: Systemic lupus erythematosus Complications: Persistent fever for 5 weeks and months postinifection. 2 months postinifection. 2 arthroplasty. 10 months postinifection acute duetrion, respiratory insufficiency and sepsis. Outcome: Death	۵
Cherabuddi et al ²⁶ (2016)	Colombia	2016	н 4	40	CHIKV	DENV	+	+	+-	+	щ	۳	۳	R	Back pain, retro-ocular pain complications : Persistent severe arthralgia after 2 months Outcome: Sequelae	ى س
Zambrano <i>et</i> a/ ²⁷ (2016)	Ecuador	2016	Σ 4	43 0	CHIKV	DENV	+-	+-	+-	+	RN	RN	+	NR	Outcome: Full recovery	5
Zambrano <i>et</i> af ²⁷ (2016)	Ecuador	2016	н 4	43 (CHIKV	RR	++	+-		+-	+	+	RN	NR	Outcome: Full recovery	CJ
Zambrano ef al ²⁷ (2016)	Ecuador	2016	с) L	57 0	CHIKV	DENV	R	+	Щ	R	RN	+	ц	R	Lumbar back pain Complications: Guillain- Barré syndrome (EMG confirmed) Outcome: Full recovery	Ŋ
Azeredo et al ^{ia} Brazil (2018)	Brazil	2016	z س	Ř	μ	OHIKA	÷	+	٣	٣	۳	+	٣	÷	Retro-orbital pain complications: Suspected vertical transmission, pregnancy outcome alive newborn with functional plagiosephaly a flat spot on the back or side of newborn's skull Gestational age at infection: 9 weeks	ю
																Continued

Table 4 C	Continued															
						Other	WHO 2	zIKV sigr	WHO ZIKV signs or symptoms	smc			Other repor symptoms	Other reported signs or symptoms		
Author (year)	Location	Study year	Sex		Age Coinfecting (years) agent(s)	tested (negative)	Rash	Fever	Arthralgia	Conjunctivitis		Myalgia Headache	URT symptoms	GI symptoms	Additional information	Level of evidence*
Azeredo et al ¹⁹ 1 (2018)	Brazil	2016	ш	R	DENC	CHIK	+-	+	щ	÷	R	۳	щ	Ч	Retro-orbital pain Complications: Suspected vertical transmission to deceased insufficiency Gestational age at infection: 20 weeks	ы
Dupont- Rouzeyrol <i>et</i> al ³² (2015)	New Caledonia	RN	ш	38	DENV 1	RN	+	+-	÷	÷	+	+	RN	÷	Asthenia, retro-ocular pain Outcome: Full recovery	Ω.
Dupont- Rouzeyrol <i>et</i> al ³² (2015)	New Caledonia	RN	Σ	14	DENV 3	RN	В	÷	+-	NR	+	÷	RN	RN	Asthenia Outcome: Full recovery	Ω
lovine <i>et al</i> ³³ (201 <i>7</i>)	United States§	2016	ш	26	DENV 2	CHIKV	+	+-	R	÷	NR	RN	+-	÷	Retro-orbital pain, fatigue, malaise, facial flushing Outcome: Full recovery	Ω
Villamil-Gomez Colombia et ar ³⁵ (2016)	Colombia	КN	ш	ĸ	CHIKV, DENV	Plasmodium spp	+	Ч И И	÷	÷	++	÷	Ψž	++	Physical examination revealed cervical lymphadenopathy, lymphadenopathy, apainful oedema and painful oedema in the lower limbs Outcome: Weekly Outcome: Weekly fow 14.6 to 29 weeks of form 14.6 to 29 weeks of gestation were normal	۵
Gunturiz <i>et al</i> ³⁶ Colombia (2018)	Colombia	2016	ш	ά	CHIKV, Toxo	R	Ë	щ	Ř	ж	RN	щ	ŭ	ж	Complications: Suspected vertical transmission of ZIKV infections to the fetus with outcome of fetus at 20 weeks of gestation, termination at 29 weeks of gestation	ۍ
Villamil-Gomez Colombia et al ⁹⁴ (2018)	Colombia	2015/2016	Σ	28	≥H	щ	+	+	щ	÷	RN	щ	Ϋ́	Ř	Recently diagnosed with HIV (<1 year), Lymphocytes T CD4 count (cells/mm ³): 450, HIV viral load (RNA copies/mL): 100 Complication: Demyelination was found (EMG findings)	ى س
																Continued

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Table 4 C	Continued															
							NHO ZII	KV signs	WHO ZIKV signs or symptoms	ns			Other reported signs or symptoms	ted signs or		
Author (year)	Location	Study year	Sex	Age (years)	ing	tested (negative) F	Rash	Fever /	Arthralgia	Conjunctivitis	Myalgia	Headache	URT symptoms	GI symptoms	Additional information	Level of evidence*
Villamil-Gomez Colombia et al ^{%4} (2018)	Colombia	2015/2016	ш	49	≥ H	۲ ۲	۳. ۳.	ч щ	۲. ۲	۳	щ	щ	Ч	+	Recently diagnosed with HIV (<1 year) Lymphocytes T CD4 count (cells/mm ³): 98 HIV viral load (RNA copies/mL): 1800 Hypotension, dysathria, decreased muscle strength, relaxation of sphincters, areflexia and basal bilateral crackles in the lungs Complications: Sepsis	۵
Villamil-Gomez Colombia et al ⁶⁴ (2018)	Colombia	2015/2016	ш	45	PH	H H		+	۲. ۲	+	Ч	Ř	Я	÷	Recently diagnosed with HIV (1 year ago) Lymphocytes T CD4 count (cells/mm ³): 380 HIV viral load (RNA copies/mL): 800 Compilication: Demyelination was found (EMG findings)	۵
Valdespino- Vazquez et al ^s (2018)	Mexico	2016	ш	22	EBV, HHV6	DENV, CHIKV, WNV, VZ, HSV-1/2, HHV7, HHV8,CMV		+	щ	٣	а И	÷	щ	R	Infection of pregnant women at 14 weeks of gestation Complications: Suspected vertical transmission of ZIKV infections to the foetus with outcome of diagnosed CZS and fetal death at 30 weeks of gestational age, 4 hours after birth	ى س
Araujo <i>et al³⁸</i> (2018)	Brazil	2016	Σ	26	HSV-1	DENV, HSV,VZ, N EBV, Toxo, HepC/B, Syphilis spp.	щ	+	Ч	RN	R	÷	RN	÷	Complications: Meningoencephalitis Outcome: Full recovery	ų
Biron et al ³⁹ (2016)	New Caledonia	R	Σ	19 1	Leptospira spp	DENV	щ	+	н К	R	÷	RN	÷	R	Complications: Haemodynamic condition was unstable, sceptic shock Outcome: Full recovery	ى
																Continued

Table 4	Continued														
					Other	MHO ZI	KV signs	WHO ZIKV signs or symptoms	SL			Other reported signs or symptoms	ed signs or		
Author (year)	Location	Study year	Ag Sex (ye	Age Coinfecting (years) agent(s)	paurogens tested (negative)	Rash	Fever	Arthralgia	Conjunctivitis		Myalgia Headache	URT symptoms	GI symptoms	Additional information	Level of evidence*
Neaterour <i>et al</i> (2017)	<i>st al</i> Puerto Rico	2016	A 48	Leptospira spp	DENV. CHIKV	٣	+	R	ц Ц	+	÷	Ϋ́Ε.	+	Complications: Severe thrombocytopenia, persistent hypotension (BP=6040), and onset of haematochezia suggestive of an acute gastrointestinal bleed. Haemodynamic instability and haemorrhagic manifestations. Cardiac arrest. Outcome: Death.	ũ
Alves et al ^{r1} (2017)	Brazil	۳	M boy	NR, a Schistosoma boy mansoni	۳	+	+	۳. ۳.	۳	щ	ц	۳	Ë	Complications: Inflammation of the right The microscopic examination of the teetis ruled out the possibility of cancer but confirmed the diagnosis of extensive loss of testicular structure and schistosome egg-induced granulomas.	υ α
URT symptoms: pharyng *All articles were rated ac Reported to be present. Reported not to be pres Fravel associated digth GHIKV, chikungunya virus Mycobacterium tuberculo	URT symptoms: pharyngitis, sore throat, "All anticles were rated according to level Pheported to be present. Freported not to be present. § Travel associated diagnosed from Halti, § Travel associated diagnosed trom Malti. CHIKV, chikungunya virus; CMV, cytome Mycobacterium tuberculosis; NR, not rej	throat, cough, ph io level of evidenc 1 Haiti. 1 ytomegalovirus; C 1 ot reported; Tox	aryngeal co ce using the CZS, conger :o, Toxoplasr	URT symptoms: pharyngits, sore throat, cough, pharyngeal congestion, adenopathy. GI symptoms: nausea, diarrhoea, vomiting, constipation, si "All articles were rated according to level of evidence using the Oxford Centre for Evidence-based Medicine's Levels of Evidence, March 2009. ¹¹ Flepported to be present. Flepported not to be present. Si Tavel associated diagnosed from Haiti. CHIKV, chikungunya virus; CMV, cytomegalovirus; CZS, congenital Zika syndrome; DENV, dengue virus; EMG, electromyography testing; GI, gas Mycobacterium tuberculosis; NR, not reportect; Toxo, Toxoplasma gondi; URT, upper respiratory tract; VZ, varicella zoster; WNV, West Nile virus.	il symptoms: nausea, o ance-based Medicine's and dengue virus: EMG espiratory tract; VZ, va	diarrhoea, vo Levels of Ev , electromyc	idence, Mi vidence, Mi ography tes r; WNV, W∈	r ausea, diarrhoea, vomiting, constipation, stomach ache. Aedicine's Levels of Evidence, March 2009. ¹¹ irus: EMG, electromyography testing: Gl, gastrointestinal, lict; VZ, varicella zoster, WNV, West Nile virus.	rach ache. intestinal; Hep, he	patitis virus	; HHV, human her	pes virus; HSV, F	erpes simplex virus	URT symptoms: pharyngits, sore throat, cough, pharyngeal congestion, adenopathy. GI symptoms: nausea, diarrhoea, vomiting, constipation, stomach ache. "All articles were rated according to level of evidence using the Oxford Centre for Evidence-based Medicine's Levels of Evidence, March 2009. ¹¹ Theopreted not be present. Free present do to be present. 6 Trave present according to level of evidence using the Oxford Centre for Evidence-based Medicine's Levels of Evidence, March 2009. ¹¹ Theopreted not to be present. 7 Free present according to level of evidence using the Oxford Centre for Evidence based Medicine's Levels of Evidence, March 2009. ¹¹ Theopreted not to be present. 6 Trave present diagnosed from Hatti. CHIKV, chikungunya virus; CMV, cytomegalovirus; CZS, congenital Zika syndrome; DENV, dengue virus; EMG, electromyography testing; GI, gastrointestinal; Hep, hepatitis virus; HHV, human herpes virus; HTV, herpes simplex virus; HTV, human herpes virus; HTV, herpes simplex virus; HTV, human T-lymphotropic virus; MTB, Mycobacterium tubercucisis; NR, not reported; Toxo, Toxoplasma gondi; URT, upper respiratory tract; VZ, varicella zoster; WNV, West Nile virus.	rus; MTB,

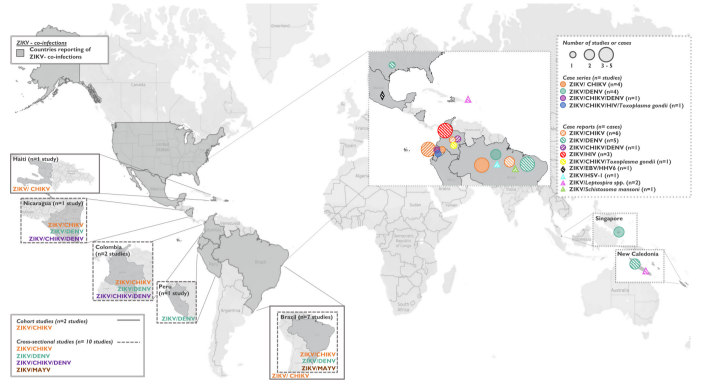


Figure 2 Studies included in the systematic review: cohort studies (n=2), cross-sectional studies (n=10), case series studies (n=8) and case reports (n=21 reported in 14 case report studies). Two cohort studies on ZIKV/CHIKV coinfections were conducted in Haiti (n=1) and Brazil (n=1). Ten cross-sectional studies were conducted in Brazil (n=6), Colombia (n=2), Nicaragua (n=1) and Peru (n=1). Eight case series were reported from Brazil (n=5), Ecuador (n=1) and Singapore (n=2). Twenty-one case reports were reported from Brazil (n=6), Colombia (n=6), Ecuador (n=3), Mexico (n=1), New Caledonia (n=3), Puerto Rico (n=1) and the USA (n=1). CHIKV, chikungunya virus; DENV, dengue virus; EBV, Epstein-Barr virus; HSV, herpes simplex virus; MAYV, Mayaro virus; ZIKV, Zika virus.

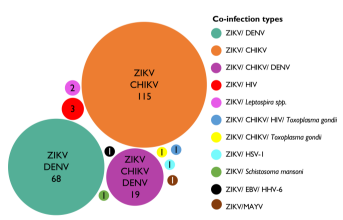


Figure 3 Zika virus coinfection types identified in this systematic review. Size of circles represents the number of cases reported per coinfection type. In total, 213 coinfection cases were included, ie, ZIKV/CHIKV (n=115), ZIKV/ DENV (n=68), ZIKV/CHIKV/DENV (n=19), ZIKV/HIV (n=3), ZIKV/Leptospira spp (n=2), ZIKV/HIV/*Toxoplasma gondii* (n=1), ZIKV/CHIKV/*Toxoplasma gondii* (n=1), ZIKV/EBV/HIV-6 (n=1), ZIKV/CHIKV/MAYV (n=1. CHIKV, chikungunya virus; DENV, dengue virus; EBV, Epstein-Barr virus; HHV, human herpes virus; HSV, herpes simplex virus; MAYV, Mayaro virus; ZIKV, Zika virus.

presented respectively with painful hepatomegaly, liver enlargement, mucosal bleeding, gingival bleeding, significant thrombocytopenia and abrupt platelet decrease.¹⁵ ¹⁸ ¹⁹ The only neurological complications resulting from ZIKV/DENV coinfection were reported in two case reports documenting infections in pregnancy, with one case resulting in a newborn with functional plagiocephaly and the other in fetal death (table 3).¹⁹

ZIKV/CHIKV/DENV coinfections

The clinical presentation of five cases with ZIKV/ CHIKV/DENV coinfection were described in one case series (four cases) and one case report (tables 3 and 4, online supplementary tables 1 and 2).^{22 34} Similar to ZIKV/CHIKV and ZIKV/DENV-coinfected cases, ZIKV/ CHIKV/DENV-coinfected cases presented with signs and symptoms consistent with the ZIKV WHO clinical case definition.¹ All five cases were reported to have complications (figure 4). The case series reported GBS in two cases, one case of meningitis and one case of encephalitis, which resulted in death. Notably, the study's population was selected to include only clinical patients presenting to hospital with neurological symptoms.²² The case report documented one case of cervical lymphadenopathy in pregnancy and full recovery.³⁵

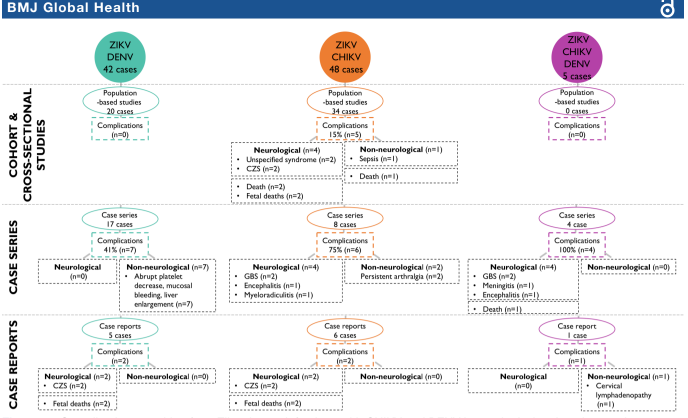


Figure 4 Complications resulting from Zika virus coinfections with CHIKV and DENV by study design. In cohort and crosssectional studies, 15% of ZIKV/CHIKV coinfections resulted in complications. In case series, 41% of ZIKV/DENV, 75% of ZIKV/ CHIKV and 100% of ZIKV/CHIKV/DENV cases resulted in in complications. In case reports, two ZIKV/DENV, two ZIKV/CHIKV and one ZIKV/CHIKV/DENV coinfections resulted in complications. CHIKV, chikungunya virus; CZS, congenital Zika syndrome; DENV, dengue virus; GBS, Guillain-Barré syndrome; n, number of complications; ZIKV, Zika virus.

Other ZIKV coinfections

There is limited published evidence on ZIKV coinfections with other pathogens. To date, the clinical signs and symptoms of 10 cases with eight other ZIKV coinfection types have been documented in one cross-sectional study, one case series and seven case reports (tables 2–4, online supplementary tables 1 and 2).^{34 36–42}

In addition to presenting with signs and symptoms consistent with the WHO ZIKV clinical case definition, almost all cases of ZIKV coinfections with pathogens other than DENV or CHIKV were reported to experience complications. Neurological complications were reported in two ZIKV/HIV coinfections, one ZIKV/ CHIKV/HIV/Toxoplasma gondii coinfection and one ZIKV/HSV-1 coinfection. These neurological complications included meningitis, meningoencephalitis and demyelinations confirmed by electromyography.^{22 34 38} Further, one ZIKV/HIV-coinfected case developed sepsis, resulting in death.³⁴ Two ZIKV/Leptospira spp-coinfected cases developed haemodynamic instability, one resulting in septic shock, and one in death.^{39 40} Additionally, one ZIKV/Schistosma mansoni-coinfected case experienced testicular inflammation with granulomas induced by schistosome eggs.⁴¹

Coinfections in pregnancy were described in three ZIKV coinfection types: ZIKV/MAYV, ZIKV/CHIKV/ *Toxoplasma gondii* and ZIKV/EBV/HHV6 coinfections.^{36 37 42} In the latter two, vertical ZIKV transmission was suspected, as both fetuses were diagnosed with CZS. After diagnosis, one pregnancy was terminated at 29 weeks of gestation and one newborn died 4 hours after birth at 30 weeks of gestation due to respiratory distress syndrome.

Levels of evidence

The levels of evidence for the studies were assessed using the OCEBM Levels of Evidence (1=highest, 5=lowest). Two cohort studies with limited follow-up were graded evidence level 2b.^{5 20} Ten cross-sectional studies were graded evidence level 2c.^{9 12-14 17 19 28 30 42 43} Eight case series studies were graded evidence level 4.^{15 16 18 21-23 29 31} Fourteen case report studies were graded evidence level 5.^{19 24-27 32-41} Thus, most of the studies included in the systematic review are evidence level 4 or 5.

DISCUSSION

This systematic review summarises the existing literature on ZIKV coinfections. Specifically, it describes the estimated frequencies of reported ZIKV coinfections and their clinical spectrum. The search identified 34 studies conducted between 2014 and 2019, which reported 213 cases of ZIKV coinfection with 10 different pathogens. ZIKV coinfections were detected across 10 countries, primarily in Latin America. CHIKV and DENV were the predominantly reported ZIKV coinfecting agents and the only ZIKV coinfections for which population frequencies were described. ZIKV coinfection frequencies among ZIKV-infected cases varied significantly between location and population type. The vast majority of ZIKV-coinfected cases were reported to present with the signs and symptoms described for uncomplicated ZIKV monoinfections and defined by the WHO.¹ However, complications were reported to arise in 9% of ZIKV-coinfected cases in cohort and cross-sectional studies.

This is the first systematic review to study how frequently individuals with ZIKV infection have a coexisting infection of any kind. The variation in frequencies reported for ZIKV/arbovirus coinfections among the ZIKV-infected individuals reported in this study was likely influenced by differences in study design and the selected study population. Factors, such as study location, season and study period in relation to the ZIKV outbreak, will have additionally influenced ZIKV coinfection frequency estimates. As expected, ZIKV coinfections were relatively more common in studies conducted during concurrent arbovirus outbreaks.^{14 44} These differences in study design, timing and location make it difficult to generalise ZIKV coinfection frequency estimates, but provide important knowledge that arbovirus coinfections can occur in up to half of ZIKV-infected cases in certain contexts. Our findings are consistent with a systematic review of CHIKV/DENV coinfections, which found the frequency of CHIKV/DENV coinfections reported in 28 studies ranged from 1% to 36%.45 The heterogeneity across studies also reflects the difficulty in estimating the background level of ZIKV infections (ie, the denominator for assessing coinfection frequencies), given the diagnostic challenges in identifying acute ZIKV infections.⁴⁶

Overall, the evidence identified in this review suggests that ZIKV coinfections appear to present with a mild clinical presentation similar to that previously described for ZIKV monoinfections. Of note, GI and URT symptoms, which are considered uncharacteristic for ZIKV, were reported to occur not infrequently in ZIKV/DENV, ZIKV/CHIKV and ZIKV/CHIKV/DENV-coinfected cases. While the evidence base from animal model studies of ZIKV coinfection is limited to date, two studies have compared ZIKV infection among rhesus macaque models with and without simian immunodeficiency virus or chimeric simian HIV.47 48 Whereas coinfected macaques were observed to have lower peak Zika viral loads with a longer clearance time in both investigations, the area under the viral load curves did not appear to differ substantively by coinfection status, potentially suggesting an overall limited impact of coinfection on disease progression but raising questions about the role of lentiviral coinfection in onward transmission.4748

Although the existing reports suggest that coinfections do not appear to markedly alter the clinical presentation of uncomplicated ZIKV disease in humans, the findings from this review highlight a need for additional high quality research investigating whether coinfections may influence complication risks. Based on the limited available evidence, the complications described for ZIKV coinfections appear to be broadly similar to those reported for ZIKV monoinfections.⁴⁹ However, 33% of the coinfection-related complications appeared to be atypical for ZIKV monoinfections, but were consistent with complications previously documented for the coinfecting pathogens (eg, bleeding in 10% of ZIKV/DENV cases and persistent arthralgia in 6% of ZIKV/CHIKV cases).^{50 51} In addition, among deaths of ZIKV-coinfected cases, three of the nine cases had immune deficiencies and one ZIKV/Leptospira spp-coinfected case died from complications established for Leptospira spp infections.⁴⁰ The remaining five deaths reported from ZIKV coinfections were three fetal deaths, one case following multiorgan failure and one case following encephalitis.^{22 28} Additionally, some complications may have been missed, especially those that occurred after the acute infections, as the follow-up period of the individual studies may have not been adequate to detect late-onset complications. Further research (eg, an ongoing cohort study of ZIKV/ HIV coinfections in pregnant women⁵²) will be valuable for discerning the relative risk of complications of ZIKV coinfection versus monoinfections.

This review had strengths and limitations. ZIKV is an emerging infectious disease of significant public health concern, and this is the first systematic review of the frequency, types and clinical presentation of ZIKV coinfections. The study employed a broad search strategy including search terms for all potential coinfecting pathogens and using multiple languages to identify all available evidence. Most importantly, the review included only qRT-PCR-confirmed ZIKV coinfections, which is the most accurate way to diagnose acute coinfections (ie, due to the very short time window of qRT-PCR testing (<7 days)) and limits misdiagnosis, which is of particular importance with the high cross-reactivity reported from arbovirus serology testing. On the other hand, by focusing on concurrent infections, the current review was unable to appraise the potential impact of recent infections; for example, it has been previously reported that pre-existing immunity to DENV, which shares a common vector and circulates in most of the countries reporting ZIKV coinfection, may influence the clinical presentation of ZIKV infection.⁵³ The additional limitations of this review mainly stem from the lack of available high-quality evidence on ZIKV coinfections. Notably, the majority of included studies were rated level 4 or 5 according to the OCEBM Levels of Evidence. Only seven studies were rated level 2 or above. Additionally, the reported ZIKV coinfection types may have been influenced by the underlying prevalence of coinfecting pathogens in the population and the applied diagnostic practices (ie, multiplex testing vs testing on clinician's suspicion). The use of specific case definitions in included cross-sectional and case series studies (eg, fever and rash¹⁵) may have also introduced a selection bias that potentially led to an overrepresentation of specific symptoms associated with ZIKV

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coinfection reported for a given study (eg, reporting 100% of cases as presenting with fever and rash).¹⁵ Finally, the studies selected for this systematic review only included symptomatic ZIKV-infected cases, which represent only approximately 40% of all ZIKV cases.² It is likely that the actual frequency of ZIKV coinfections may be higher as many cases will be asymptomatic and therefore never seek medical attention. However, the recently implemented multiplex PCR assay, which tests for CHIKV, DENV and ZIKV simultaneously, will likely improve the detection of ZIKV/arbovirus coinfections and facilitate future assessment of the frequency of ZIKV coinfections.⁵⁴

In conclusion, the findings of this review suggest that the cocirculating arboviruses, CHIKV and DENV, are the most common ZIKV coinfection types and may, in specific populations and epidemiological contexts, occur in up to half of ZIKV infections. The evidence collated in this systematic review suggests coinfections do not markedly alter the generally mild clinical presentation of uncomplicated ZIKV disease. However, additional and better quality evidence should be prioritised in future outbreaks to corroborate the estimates of the frequency of ZIKV coinfections and to interrogate the importance of ZIKV coinfections in the development of ZIKV-related complications, especially for ZIKV coinfections with CHIKV and DENV.

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Contributors All authors contributed substantially to the design of the work and/ or the acquisition, analysis and interpretation of the data, contributed meaningfully to the drafting and/or revision of the manuscript, provided final approval for the version published and share responsibility for the published findings.

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Competing interests None declared.

Patient and public involvement Patients and/or the public were not involved in the design, or conduct, or reporting or dissemination plans of this research.

Patient consent for publication Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement All data relevant to the study are included in the article or uploaded as supplementary information. This systematic review is based on published articles. All abstracted data are provided in the text and supplementary materials.

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